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Case report

Acute infarction of corpus callosum due to transient obstructive hydrocephalus



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AND NEUROSURGERY

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ABSTRACT

Acute ischemia of the corpus callosum (CC) is not a well-known feature in patients with acute hydrocephalus. Herein, we describe a case with acute CC infarction due to another rare entity; transient obstructive hydrocephalus. A 66-year-old male was admitted with sudden onset right-sided hemiparesia. CT demonstrated a hematoma on the left basal ganglia with extension to all ventricles. The following day, the patient's neurological status progressed to coma and developed bilateral pyramidal signs. MRI demonstrated obstructive hydrocephalus and acute diffuse infarction accompanied by elevation of the CC. On the same day there was improvement in his neurological status with significant decrease in ventricular size and complete resolution of the clot in the third ventricle. The mechanism of signal abnormalities is probably related with the neural compression of the CC against the falx. Presumably, the clot causing obstruction in the third ventricle dissolved or decayed by the help of fibrinolytic activity of CSF, which was raised after IVH and caused spontaneous improvement of hydrocephalus. Bilateral neurological symptoms suggest diffuse axonal damage and normalization of the intracranial pressure should be performed on the early onset of clinical detorioration in order to prevent axonal injury.

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1. Introduction

Arterial supply of corpus callosum (CC) is derived from both anterior and posterior circulation.

Therefore, infarcts of CC are not common and mainly caused by infarctions of anterior cerebral artery and posterior cerebral artery territories. Because of the rich blood supply, solitary infarct of the CC is rare and generally associated with infarcts of the neighboring structures [1]. Morphological changes such as elevation, thinning and impingement of CC have previously been described in acute hydrocephalus whereas acute ischemia is not a well-known feature [2–4]. Herein, we describe a case with acute CC infarction due to another rare entity; transient posthemorrhagic obstructive hydrocephalus.

2. Case report

A 66-year-old male was admitted to our hospital with the chief complain of sudden onset of right sided hemiparesia. He had hypertension in his past medical history. On neurological examination, he was alert with full cooperation and orientation,

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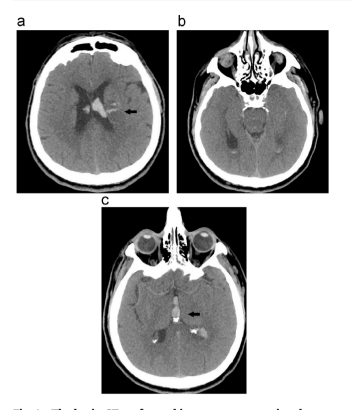
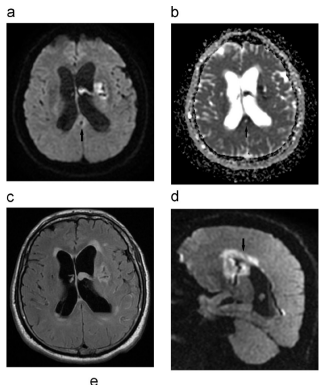


Fig. 1 – The brain CT performed in emergency service shows a left sided basal ganglia hematoma (a) with extension to all ventricles (b) and the clot in the third ventricle (c).

he had right sided central facial palsy, hemiparesia, hemihypoesthesia and the Babinski sign. The computerized tomography (CT) of the brain showed a left sided basal ganglia hematoma with extension to all ventricles, the volume of intraventricular hemorrhage was 9 ml (Fig. 1). The patient was hospitalized and followed in the intensive care unit. Twentytwo hours after presentation, he became stuporous and CT showed ventricular enlargement. Four hours later he underwent brain magnetic resonance imaging (MRI) to exclude other reasons of clinical impairment. MRI demonstrated acute obstructive hydrocephalus, elevation and hyperacute infarction of the corpus callosum (Fig. 2). He was intubated; hyperventilation and mannitol therapy was started. The neurological status progressed to coma within hours and he developed extensor response to noxious stimuli and bilateral Babinski sign. Although there was no cause explaining neurological impairment other than hydrocephalus, the neurosurgery department recommended following the patient with medical therapy and did not perform any intervention. Sixteen hours later, neurological deterioration suddenly ceased and later on he was able to localize painful stimuli. The CT obtained the next day, 48 h after initial presentation showed significant decrease in the ventricular size and complete resolution of the clot in the third ventricle (Fig. 3). The CC infarction was more prominent on the MRI which was obtained 12 days after the incident (Fig. 4). Digital subtraction angiography did not reveal any vascular pathology. Consequently, the etiology of the bleeding was considered as hypertensive hemorrhage. He was discharged two months later and was able to cooperate to all simple and some complex



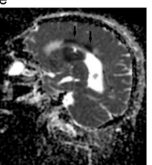


Fig. 2 – Axial diffusion (a), apparent diffusion coefficient (ADC) (b), sagittal diffusion (d) sagittal ADC (e) sequences of the brain MRI demonstrate hemorrhage in left basal ganglia and the infarction of the CC. The fluid-attenuated inversion recovery (FLAIR) sequence does not show any signal changes (c) in CC indicating hyperacute phase of ischemia. Elevation of the CC was seen in the sagittal sequences.

commands and his muscle strength had recovered to grade 4/5 in all four limbs.

3. Discussion

We described a recent case of acute infarction of the CC due to acute obstructive hydrocephalus. The subcallosal and medial callosal arteries supply blood to the anterior part of the CC, the pericallosal artery supplies the body, while the posterior pericallosal artery is responsible for the blood supply of the splenium. Because of its rich blood supply, the CC is known as a resistant area for ischemia. Hence isolated callosal infarcts Download English Version:

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